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NEWS	1			Web Page for STN Seminar Schedule - N. America
NEWS	2	JAN	02	STN pricing information for 2008 now available
NEWS	3	JAN	16	CAS patent coverage enhanced to include exemplified
				prophetic substances
NEWS	4	JAN	28	USPATFULL, USPAT2, and USPATOLD enhanced with new
				custom IPC display formats
NEWS	5	JAN	28	MARPAT searching enhanced
NEWS				USGENE now provides USPTO sequence data within 3 days
				of publication
NEWS	7	JAN	28	TOXCENTER enhanced with reloaded MEDLINE segment
NEWS		JAN		MEDLINE and LMEDLINE reloaded with enhancements
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		FEB		IMSPRODUCT reloaded with enhancements
NEWS	13	FEB	29	WPINDEX/WPIDS/WPIX enhanced with ECLA and current
				U.S. National Patent Classification
NEWS	14	MAR	31	IFICDB, IFIPAT, and IFIUDB enhanced with new custom
				IPC display formats
NEWS	15	MAR	31	CAS REGISTRY enhanced with additional experimental
				spectra
NEWS	16	MAR	31	CA/CAplus and CASREACT patent number format for U.S.
				applications updated
NEWS	17	MAR	31	LPCI now available as a replacement to LDPCI
NEWS	18	MAR	31	EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS	19	APR	0.4	STN AnaVist, Version 1, to be discontinued
		APR		WPIDS, WPINDEX, and WPIX enhanced with new
				predefined hit display formats
NEWS	21	APR	28	EMBASE Controlled Term thesaurus enhanced
NEWS	22	APR	28	IMSRESEARCH reloaded with enhancements
		MAY		INPAFAMDB now available on STN for patent family
				searching
NEWS	24	MAY	3.0	DGENE, PCTGEN, and USGENE enhanced with new homology
112110				sequence search option
NEWS	25	JUN	06	EPFULL enhanced with 260,000 English abstracts
NEWS				KOREAPAT updated with 41,000 documents
112112		0011	0.0	normalita apadeed water 12,000 documentes
NEWS	EXE	RESS	FEB	RUARY 08 CURRENT WINDOWS VERSION IS V8.3,
				CURRENT DISCOVER FILE IS DATED 20 FEBRUARY 2008
NEWS	HOL	IRS	ST	N Operating Hours Plus Help Desk Availability
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SINCE FILE

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TOTAL ENTRY SESSION

0.63

FILE 'HOME' ENTERED AT 23:16:40 ON 08 JUN 2008

=> set abbr on perm SET COMMAND COMPLETED

=> set plurals on perm SET COMMAND COMPLETED

=> file uspatall caplus japio COST IN U.S. DOLLARS

FULL ESTIMATED COST

FILE 'USPATFULL' ENTERED AT 23:18:08 ON 08 JUN 2008 CA INDEXING COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'USPATOLD' ENTERED AT 23:18:08 ON 08 JUN 2008 CA INDEXING COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'USPAT2' ENTERED AT 23:18:08 ON 08 JUN 2008 CA INDEXING COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'CAPLUS' ENTERED AT 23:18:08 ON 08 JUN 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'JAPIO' ENTERED AT 23:18:08 ON 08 JUN 2008 COPYRIGHT (C) 2008 Japanese Patent Office (JPO) - JAPIO

=> s (imidat? or imidiz? or imidis?) and supercritical? 159 (IMIDAT? OR IMIDIZ? OR IMIDIS?) AND SUPERCRITICAL?

=> s sma (s)(polymer? or copolymer?) L2 4630 SMA (S) (POLYMER? OR COPOLYMER?)

=> s 11 and 12 L3 0 L1 AND L2

=> s (styren? or vinv1(2a)aromatic)(3a)copoly? or poly(1w)styren?(1w)co? 1 FILES SEARCHED...

TERM 'CO?' EXCEEDED TRUNCATION LIMITS - SEARCH ENDED You have entered a truncated stem which occurs in too many terms. Make the stem longer and try again. For example, if your original term was 'degr?' to search for variations and the abbreviation for 'degradation', you could replace it with the expression '(degrdn OR degrad?)'. If your search term was numeric, e.g., 'C>5', reduce the size of the range.

=> s (styren? or viny1(2a)aromatic)(3a)copoly? or poly(1w)styren?(3w)maleic

279533 (STYREN? OR VINYL(2A) AROMATIC)(3A) COPOLY? OR POLY(1W) STYREN?(3W) MALEIC ANHYDRIDE

=> s 11 and 14 11 L1 AND L4

=> d 15 1-11 ibib abs

L5 ANSWER 1 OF 11 USPATFULL on STN

ACCESSION NUMBER: 2008:118581 USPATFULL

TITLE: Ink-jet ink and cured film obtained from same

INVENTOR(S): Satou, Hiroyuki, Ichihara-shi, JAPAN Deyama, Yoshihiro, Ichihara-shi, JAPAN

Yamahiro, Mikio, Ichihara-shi, JAPAN PATENT ASSIGNEE(S): CHISSO Corporation (non-U.S. corporation)

KIND DATE NUMBER ______ PATENT INFORMATION: US 20080103280 A1 20080501 US 2007-976689 A1 20071026 (11) APPLICATION INFO.:

> NUMBER DATE

_____ PRIORITY INFORMATION: JP 2006-290740 20061026

DOCUMENT TYPE: Utility

APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: HOGAN & HARTSON LLP, IP GROUP, COLUMBIA SQUARE, 555 THIRTEENTH STREET, N.W., WASHINGTON, DC, 20004, US

NUMBER OF CLAIMS: 33

EXEMPLARY CLAIM:

LINE COUNT: 2088 CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention provides an ink-jet ink including a fluorine-containing compound (C) in the form of fluorosilsesquioxane having an organic group having 1 to 100 carbon atoms.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 2 OF 11 USPATFULL on STN

ACCESSION NUMBER: 2007:140924 USPATFULL

TITLE: Continuous process for the production of combinatorial

libraries of modified materials Nelson, James M., Roseville, MN, UNITED STATES INVENTOR(S):

Marx, Ryan E., Cottage Grove, MN, UNITED STATES Cernohous, Jeffrey J., Hudson, WI, UNITED STATES

McNerney, James R., Inver Grove Heights, MN, UNITED

PATENT ASSIGNEE(S): 3M Innovative Propeties Company (U.S. corporation)

NUMBER KIND DATE -----PATENT INFORMATION: US 20070122915 A1 20070531 APPLICATION INFO.: US 2006-646856 A1 20061228 (11)

RELATED APPLN. INFO.: Continuation of Ser. No. US 2002-211219, filed on 2 Aug

2002, GRANTED, Pat. No. US 7157283 Utility

DOCUMENT TYPE: APPLICATION

LEGAL REPRESENTATIVE: 3M INNOVATIVE PROPERTIES COMPANY, PO BOX 33427, ST.

PAUL, MN, 55133-3427, US

NUMBER OF CLAIMS: 20 EXEMPLARY CLAIM: 1

LINE COUNT: 1325

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A system is provided wherein a devolatilizing reactor is used to make combinatorial libraries of materials. Examples of suitable reactors include continuous high viscosity devolatilizers and continuous devolatilizing kneaders.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 3 OF 11 USPATFULL on STN

ACCESSION NUMBER: 2007:107637 USPATFULL

TITLE: Method for preparing styrene and maleimide copolymer using super critical fluid

Kim, Dong Ryul, Daejeon, KOREA, REPUBLIC OF INVENTOR(S): Park, Sang Hyun, Daejeon, KOREA, REPUBLIC OF

Lee, Hee Hyun, Daejeon, KOREA, REPUBLIC OF

NUMBER KIND DATE PATENT INFORMATION: APPLICATION INFO.: US 20070093610 A1 20070426 US 2004-554708 A1 20040512 (10) WO 2004-KR1090 20040512 20051027 PCT 371 date

DOCUMENT TYPE: Utility FILE SEGMENT:

APPLICATION LEGAL REPRESENTATIVE: Mckenna Long & Aldridge LLP, 1900 K Street, NW,

Washington, DC, 20006, US

29 NUMBER OF CLAIMS:

EXEMPLARY CLAIM: LINE COUNT: 593

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Disclosed is a method for preparing copolymers of styrene and maleimide by imidization-extrusion of copolymers of styrene and maleic anhydride using a

supercritical fluid. The method can be performed at a lower

temperature compared to a conventional method and can remove residual amines and by-products with ease, and thus can provide

copolymers of styrene and maleimide having excellent

optical properties, thermal properties and mechanical properties.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 4 OF 11 USPATFULL on STN

2004:231455 USPATFULL ACCESSION NUMBER:

TITLE: Process of manufacturing optical waveguide and

connection structure of optical devices

INVENTOR(S): Oe, Kunishige, Kyoto, JAPAN

Yamashita, Kenichi, Kvoto, JAPAN Mune, Kazunori, Osaka, JAPAN Mochizuki, Amane, Osaka, JAPAN Naitou, Ryuusuke, Osaka, JAPAN

PATENT ASSIGNEE(S): NITTO DENKO CORPORATION (U.S. individual)

NUMBER KIND DATE PATENT INFORMATION: US 20040178522 A1 20040916 APPLICATION INFO.: US 2004-760389 A1 20040121 (10)

NUMBER DATE PRIORITY INFORMATION: JP 2003-15533 20030124 DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION LEGAL REPRESENTATIVE: SUGHRUE MION, PLLC, 2100 PENNSYLVANIA AVENUE, N.W.,

SUITE 800, WASHINGTON, DC, 20037

NUMBER OF CLAIMS: 12

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 5 Drawing Page(s) LINE COUNT: 928

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention provides a process of manufacturing an optical waveguide for optically connecting a plurality of optical devices, comprising the steps of: disposing a resin composition between two or more optical devices, the resin composition comprising a resin and a 1,4-dihydropyridine derivative, forming an optical path through the resin composition between the optical devices by light having a wavelength capable of inducing a structural change in the 1,4-dihydropyridine derivative, and removing the 1,4-dihydropyridine derivative from the resulting resin composition. Also disclosed is a

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 5 OF 11 USPATFULL on STN

ACCESSION NUMBER: 2004:31988 USPATFULL

TITLE: Process to modify polymeric materials and resulting

compositions

INVENTOR(S): Nelson, James M., Roseville, MN, UNITED STATES

Marx, Rvan E., Cottaga Grove, MN, UNITED STATES Cernohous, Jeffrey J., Hudson, WI, UNITED STATES McNerney, James R., Inver Grove Heights, MN, UNITED

STATES

connection structure obtained by the process.

Jones, Todd D., St. Paul, MN, UNITED STATES Hanley, Kenneth J., Eagan, MN, UNITED STATES

NUMBER KIND DATE PATENT INFORMATION:

US 20040024130 A1 20040205 US 2002-211415 A1 20020802 (10) APPLICATION INFO.: APPLICATION
DOCUMENT TYPE: Utility
APPLICATION

LEGAL REPRESENTATIVE: 3M INNOVATIVE PROPERTIES COMPANY, PO BOX 33427, ST.

PAUL, MN, 55133-3427

NUMBER OF CLAIMS: 39

EXEMPLARY CLAIM: 1

1799 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Disclosed is a method for modifying a polymer by carrying out a thermally-induced reaction in a mixing apparatus having a high shear environment and devolatilization capabilities. Also disclosed are the resulting materials.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 6 OF 11 USPATFULL on STN

ACCESSION NUMBER: 2004:31259 USPATFULL

TITLE: Continuous process for the production of combinatorial INVENTOR(S):

libraries of modified materials

Nelson, James M., Roseville, MN, UNITED STATES Marx, Ryan E., Cottage Grove, MN, UNITED STATES Cernohous, Jeffrey J., Hudson, WI, UNITED STATES McNerney, James R., Inver Grove Heights, MN, UNITED

STATES

NUMBER KIND DATE ______ US 20040023398 A1 20040205 US 7157283 B2 20070102 PATENT INFORMATION: APPLICATION INFO.: US 2002-211219 A1 20020802 (10) DOCUMENT TYPE: Utility APPLICATION

FILE SEGMENT:

LEGAL REPRESENTATIVE: 3M INNOVATIVE PROPERTIES COMPANY, PO BOX 33427, ST.

PAUL, MN, 55133-3427

NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT: 1342

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A system is provided wherein a devolatilizing reactor is used to make combinatorial libraries of materials. Examples of suitable reactors include continuous high viscosity devolatilizers and continuous devolatilizing kneaders.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 7 OF 11 USPATFULL on STN

ACCESSION NUMBER: 2002:337815 USPATFULL

TITLE: Golf ball compositions including microcellular materials and methods for making same

Harris, Kevin M., New Bedford, MA, UNITED STATES INVENTOR(S): Rajagopalan, Murali, South Dartmouth, MA, UNITED STATES

Cavallaro, Christopher, Lakeville, MA, UNITED STATES

NUMBER KIND DATE PATENT INFORMATION: US 20020193179 A1 20021219 US 6676866 B2 20040113 US 2002-142168 A1 20020510 (10) APPLICATION INFO.:

RELATED APPLN. INFO.: Division of Ser. No. US 2000-565108, filed on 4 May 2000, GRANTED, Pat. No. US 6386992

DOCUMENT TYPE: Utility

APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: SWIDLER BERLIN SHEREFF FRIEDMAN, LLP, 3000 K STREET,

NW, BOX IP, WASHINGTON, DC, 20007

24 NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 1 Drawing Page(s)

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention is directed to golf balls including one or more foamed. microcellular materials. The invention also encompasses methods of controlling or adjusting one or more material properties or the weight distribution of a golf ball, and methods of forming golf balls including such microcellular materials.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 8 OF 11 USPATFULL on STN

ACCESSION NUMBER: 2002:108265 USPATFULL

Golf ball compositions including microcellular TITLE:

materials and methods for making same

INVENTOR(S): Harris, Kevin M., New Bedford, MA, United States

Rajagopalan, Murali, South Dartmouth, MA, United States Cavallaro, Christopher, Lakeville, MA, United States

PATENT ASSIGNEE(S): Acushnet Company, Fairhaven, MA, United States (U.S.

corporation)

NUMBER KIND DATE - --US 6386992 B1 20020514 US 2000-565108 20000504 (9) PATENT INFORMATION: APPLICATION INFO.:

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED

PRIMARY EXAMINER: Sewell, Paul T.
ASSISTANT EXAMINER: Hunter, Jr., Alvin A.

LEGAL REPRESENTATIVE: Swidler Berlin Shereff Friedman, LLP

NUMBER OF CLAIMS: 14

EXEMPLARI CLAMINGS: 3 D: 963 3 Drawing Figure(s); 1 Drawing Page(s)

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention is directed to golf balls including one or more foamed, microcellular materials. The invention also encompasses methods of controlling or adjusting one or more material properties or the weight distribution of a golf ball, and methods of forming golf balls including such microcellular materials.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 9 OF 11 USPAT2 on STN

ACCESSION NUMBER: 2004:31259 USPAT2

TITLE: Continuous process for the production of combinatorial

libraries of modified materials INVENTOR(S):

Nelson, James M., Roseville, MN, UNITED STATES Marx, Ryan E., Cottage Grove, MN, UNITED STATES Cernohous, Jeffrey J., Hudson, WI, UNITED STATES McNerney, James R., Inver Grove Heights, MN, UNITED

STATES

PATENT ASSIGNEE(S): 3M Innovative Properties Company, St. Paul, MN, UNITED

STATES (U.S. corporation)

NUMBER KIND DATE PATENT INFORMATION: US 7157283 B2 20070102 APPLICATION INFO.: US 2002-211219 20020802 (10) APPLICATION INFO.:

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED FILE SEGMENT: GRANTED
PRIMARY EXAMINER: Warden, Jill
ASSISTANT EXAMINER: Levkovich, Natalia
LEGAL REPRESENTATIVE: Edman, Sean J.

NUMBER OF CLAIMS: 25 EXEMPLARY CLAIM:

LINE COUNT: 1396

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A system is provided wherein a devolatilizing reactor is used to make combinatorial libraries of materials. Examples of suitable reactors include continuous high viscosity devolatilizers and continuous devolatilizing kneaders.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 10 OF 11 USPAT2 on STN

ACCESSION NUMBER: 2002:337815 USPAT2

TITLE: Golf ball compositions including microcellular

materials and methods for making same

INVENTOR(S): Harris, Kevin M., New Bedford, MA, United States

Rajagopalan, Murali, South Dartmouth, MA, United States Cavallaro, Christopher, Lakeville, MA, United States

PATENT ASSIGNEE(S): Acushnet Company, Fairhaven, MA, United States (U.S.

corporation)

NUMBER KIND DATE PATENT INFORMATION: US 6676866 B2 20040113 US 2002-142168 20020510 (10) APPLICATION INFO.:

RELATED APPLN. INFO .: Division of Ser. No. US 2000-565108, filed on 4 May

2000, now patented, Pat. No. US 6386992

DOCUMENT TYPE: Utility

FILE SEGMENT: GRANTED PRIMARY EXAMINER: Kuhns, Allan R.

LEGAL REPRESENTATIVE: Swidler Berlin Shereff Friedman, LLP

NUMBER OF CLAIMS: 16

EXEMPLARY CLAIM: NUMBER OF DRAWINGS: 3 Drawing Figure(s); 1 Drawing Page(s)

LINE COUNT: 973

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention is directed to golf balls including one or more foamed, microcellular materials. The invention also encompasses methods of

controlling or adjusting one or more material properties or the weight distribution of a golf ball, and methods of forming golf balls including such microcellular materials.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 11 OF 11 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:1016086 CAPLUS

DOCUMENT NUMBER: 141:424600

TITLE: Method for preparing styrene and maleimide copolymer using supercritical fluids

Kim, Dong-Ryul; Park, Sang-Hyun; Lee, Hee-Hyun INVENTOR(S):

LG Chem, Ltd., S. Korea PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 22 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
VO 2004101636	A2 20041125 A3 20050217	WO 2004-KR1090	20040512
W: AE, AG, AL	, AM, AT, AU, AZ,	BA, BB, BG, BR, BW, DM, DZ, EC, EE, EG,	
GE, GH, GM	M, HR, HU, ID, IL,	IN, IS, JP, KE, KG,	KP, KZ, LC, LK,
		MG, MK, MN, MW, MX, RU, SC, SD, SE, SG,	

FILE SEGMENT:

NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

LINE COUNT:

LEGAL REPRESENTATIVE:

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RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
             EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE,
             SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
             SN. TD. TG
     KR 2004098896
                                20041126 KR 2003-31214
                                                                    20030516
     EP 1625170
EP 1625170
                         A2 20060215 EP 2004-732516
B1 20070704
                                                                    20040512
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK
                  A 20060531 CN 2004-80011161
     JP 200522171 T 20060928 JP 2005-500694
AT 366265 T 20070715 AT 2004-732516
TW 263649 B 20061011 TW 2004-93113608
US 2007003610 A1 20070426 US 2005-554708
                                                                    20040512
                                                                    20040512
                                                                    20040514
                                                                    20051027
PRIORITY APPLN. INFO.:
                                            KR 2003-31214
                                                                A 20030516
                                             WO 2004-KR1090
                                                                W 20040512
    Disclosed is a method for preparing copolymers of styrene
     and maleimide by imidization-extrusion of copolymers
     of styrene and maleic anhydride using a supercrit.
     fluid. The method can be performed at a lower temperature compared to a
     conventional method and can remove residual amines and byproducts with
     ease, and thus can provide copolymers of styrene and
     maleimide having excellent optical properties, thermal properties and
     mech. properties.
=> d 15 5 ibib hit
L5 ANSWER 5 OF 11 USPATFULL on STN
ACCESSION NUMBER:
                        2004:31988 USPATFULL
TITLE:
                        Process to modify polymeric materials and resulting
                        compositions
INVENTOR(S):
                        Nelson, James M., Roseville, MN, UNITED STATES
                        Marx, Ryan E., Cottaga Grove, MN, UNITED STATES
                        Cernohous, Jeffrey J., Hudson, WI, UNITED STATES
                        McNerney, James R., Inver Grove Heights, MN, UNITED
                        STATES
                        Jones, Todd D., St. Paul, MN, UNITED STATES
                        Hanley, Kenneth J., Eagan, MN, UNITED STATES
                             NUMBER KIND DATE
PATENT INFORMATION:
                        US 20040024130 A1 20040205
APPLICATION INFO.:
                        US 2002-211415
                                          A1 20020802 (10)
DOCUMENT TYPE:
                        Utility
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CAS INDEXING IS AVAILABLE FOR THIS PATENT.

DETD [0046] One aspect of the present invention employs thermally-induced reactions to modify polymeric materials. Many types of thermally-induced reactions are suitable for the present invention. One suitable type of reaction is a rearrangement reaction in which the substituents or moieties of a molecule are rearranged to form a new molecule, i.e., the bonding site of a substituent or moiety moves from one atom to another in the same molecule. Another suitable type of reaction is an

3M INNOVATIVE PROPERTIES COMPANY, PO BOX 33427, ST.

APPLICATION

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1799

PAUL, MN, 55133-3427

DETD

elimination reaction in which one or more substituents is removed from a molecule. Specific types of reactions that can be carried out include, but are not limited to, pyrolysis reactions, acid-catalyzed reactions, deprotection reactions, condensation reactions, hydrolysis reactions, imidization reactions, base-catalyzed reactions, and deesterification, e.g., deacetylation. In a pyrolysis reaction, a complex molecule is broken into simpler units by the use of heat. In an acid-catalyzed reaction, acid is used to drive or induce the thermal reaction. In a deprotection reaction, a protecting group is removed to expose a reactive functional group. In a condensation reaction, two molecules react to form a new molecule and release a byproduct, which is typically water. In a hydrolysis reaction, water reacts with another molecule (e.g., ester) to form one or more new molecules. In an imidization reaction, anhydrides react with primary amines via an intermediate amic acid functionality to form an imide ring and water. In a base-catalyzed reaction, base is used to drive or induce the thermal reaction. In a deesterification reaction, an ester is converted into a carboxylic acid or an anhydride. In a deacetylation reaction, an ester is converted into an alcohol with removal of an acetyl group. See, for example, Hawker et al., Macromolecules, 1998, 31, 1024. [0066] The starting polymer systems may be synthesized in processes that are carried out in batch, semibatch, continuous stirred tank reactor (CSTR), tubular reactors, stirred tubular reactors, plug flow reactors (PFR), temperature controlled stirred tubular reactors as described in WO 0158962 Al and co-pending U.S. patent app. Ser. No. 09/824,330, static mixers, continuous loop reactor, extruders, shrouded extruders as described in WO 9740929, and pouched reactors as described in WO 9607522 and WO 9607674. The media in which the polymerizations may take place are bulk, solution, suspension, emulsion, ionic liquids or

dioxide. DETD [0075] where represents a bond in a polymerizable or polymer chain; R.sub.f is -- C.sub.6F.sub.13, -- C.sub.4F.sub.9, or -- C.sub.3F.sub.7; R and R.sub.2 are each independently hydrogen or alkyl of 1 to 20 carbon atoms; n is an integer from 2 to 11; and x is an integer of at least 1. An example of a Formula I structure is N-methylperfluorobutanesulfonamid o. The fluorinated materials may be end-functionalized at one or both terminus with reactive end groups. If there are two reactive end groups, they may be the same or different. Fluorinated diene, methacrylate and styrenic homo and block copolymers end-functionalized with alcohol(s), thiol(s), and/or amine(s) may be synthesized anionically by the use of suitable anionic initiators which contain protected functional groups that can be removed by post polymerization techniques. Suitable functional initiators are known in the art and are described in, e.g., U.S. Pat. No. 6,197,891, U.S. Pat. No. 6,160,054, U.S. Pat. No. 6,221,991, and U.S. Pat. No. 6,184,338. [0081] Various reactions may be carried out to produce DETD

supercritical fluids, such as supercritical carbon

[0081] Various reactions may be carried out to produce acetylene-containing polymers. In these reactions, a sulfoxide is pyrolyzed to give the polyactylene and a sulfenic acid byproduct (RSOH) as shown in Formula II. For example, a benzenesulfenic acid may be eliminated from poly(phenyl vinyl sulfoxide)-containing copolymers to produce polyacetylene-containing copolymers, such as poly(styrene-acetylene) block copolymers. Polyacetylene is typically difficult to work with because it is very insoluble in other materials. However, including it in a block structure allows the final structure to remain soluble.

DETD [0094] Diene, methacrylate and styrenic homo and block copolymers end-functionalized with alcohol(s), thiol(s), and/or amine(s) may be synthesized anionically by the use of suitable anionic initiators which contain protected functional groups that can be removed

by post polymerization techniques. Suitable functional initiators are known in the art and are described in, e.g., U.S. Pat. No. 6,197,891, U.S. Pat. No. 6,160,054, U.S. Pat. No. 6,221,991, and U.S. Pat. No. 6.184.338.

- [0106] The polymeric materials produced by acid-catalyzed pyrolysis of DETD methacrylic and acrylic esters are methacrylic/acrylic acid or methacrvlic/acrvlic anhydride functionalized polymers. These acid- and anhydride-functionalized polymers may participate in further grafting reactions including esterification, amidation, and imidization reactions.
- DETD [0108] In the case of amidation or imidization, the acid- or anhydride-functionalized polymeric material is subjected to reaction with amine nucleophiles. Suitable amines that participate in this reaction consist of, but are not limited to, typically primary alkyl, aryl, and alkylaryl-amines. The primary amines formula is RNH.sub.2 wherein R stands for an alkyl group having 1 to 10 carbon atoms, a cycloalkyl group having 5 to 12 carbon atoms, an aralkyl group having 7 to 22 carbon atoms or an aryl group having 6 to 12 carbon atoms.
- DETD [0110] Diene, methacrylate and styrenic homo and block copolymers end-functionalized with alcohol(s), thiol(s), and/or amine(s) may be synthesized anionically by the use of suitable anionic initiators, which contain protected functional groups that can be removed by post polymerization techniques. Suitable anionic initiators are known in the art and are described in, e.g., U.S. Pat. No. 6,197,891, U.S. Pat. No. 6,160,054, U.S. Pat. No. 6,221,991, and U.S. Pat. No. 6,184,338.
- DETD Batch Synthesis of poly(styrene-acetylene) Block Copolymers via the Thermal Modification of poly(styrene-vinyl phenyl sulfoxide).
- [0144] This example illustrates the thermal elimination of DETD benzenesulfenic acid from poly(styrene-vinyl phenyl sulfoxide) (P(S-VPS)) to produce a poly(styrene-acetylene) block copolymer.

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ANSWER 6 OF 11 USPATFULL on STN

ACCESSION NUMBER: 2004:31259 USPATFULL

TITLE: Continuous process for the production of combinatorial

libraries of modified materials

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STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 20040023398	A1	20040205	
	US 7157283	B2	20070102	
APPLICATION INFO.:	US 2002-211219	A1	20020802	(10)
DOCUMENT TYPE:	Utility			
FILE SEGMENT:	APPLICATION			
LEGAL REPRESENTATIVE:	3M INNOVATIVE PRO	OPERTIE:	S COMPANY.	PO BOX 33427, ST.
	PAUL, MN, 55133-3	3427		
NUMBER OF CLAIMS:	27			
EXEMPLARY CLAIM:	1			

LINE COUNT: 1342

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- STIMM [0063] One aspect of the present invention employs thermally-induced reactions to modify polymeric materials. Many types of thermally-induced reactions are suitable for the present invention. One suitable type of reaction is a rearrangement reaction in which the substituents or moieties of a molecule are rearranged to form a new molecule, i.e., the bonding site of a substituent or moiety moves from one atom to another in the same molecule. Another suitable type of reaction is an elimination reaction in which one or more substituents is removed from a molecule. Specific types of reactions that can be carried out include, but are not limited to, pyrolysis reactions, acid-catalyzed reactions, deprotection reactions, condensation reactions, hydrolysis reactions, imidization reactions, base-catalyzed reactions, and deesterification, e.g., deacetylation. In a pyrolysis reaction, a complex molecule is broken into simpler units by the use of heat. In an acid-catalyzed reaction, acid is used to drive or induce the thermal reaction. In a deprotection reaction, a protecting group is removed to expose a reactive functional group. In a condensation reaction, two molecules react to form a new molecule and release a byproduct, which is typically water. In a hydrolysis reaction, water reacts with another molecule (e.g., ester) to form one or more new molecules. In an imidization reaction, anhydrides react with primary amines via an intermediate amic acid functionality to form an imide ring and water. In a base-catalyzed reaction, base is used to drive or induce the thermal reaction. In a deesterification reaction, an ester is converted into a carboxylic acid or an anhydride. In a deacetylation reaction, an ester is converted into an alcohol with removal of an acetyl group. See, for example, Hawker et al., Macromolecules, 1998, 31, 1024.
- SUMM [0080] The starting polymer systems may be synthesized in processes that are carried out in batch, semibatch, continuous stirred tank reactor (CSTR), tubular reactors, stirred tubular reactors, plug flow reactors (PFR), temperature controlled stirred tubular reactors as described in W0 0158962 Al and co-pending U.S. patent app. Ser. No. 09/824,330, static mixers, continuous loop reactor, extruders, shrouded extruders as described in W0 9740929, and pouched reactors as described in W0 9607522 and W0 9607674. The media in which the polymerizations may take place are bulk, solution, suspension, emulsion, ionic liquids or supercritical fluids, such as supercritical carbon dioxide.
- SUMM [0090] represents a bond in a polymerizable or polymer chain; R.sub.f is --C.sub.6F.sub.13, --C.sub.4F.sub.7, or --C.sub.5F.sub.7, R and R.sub.2 are each independently hydrogen or alkyl of 1 to 20 carbon atoms; n is an integer from 2 to 11; and x is an integer of at least 1. The fluorinated materials may be end-functionalized at one or both terminus with reactive end groups. If there are two reactive end groups, they may be the same or different. Fluorinated diene, methacrylate and styrenic homo and block copolymers end-functionalized with alcohol(s), thiol(s), and/or amine(s) may be synthesized anionically by the use of suitable anionic initiators which contain protected functional groups that can be removed by post polymerization techniques. Suitable functional initiators are known in the art and are described in, e.g., U.S. Pat. No. 6,197,891, U.S. Pat. No. 6,160,054, U.S. Pat. No. 6, 221,991, and U.S. Pat. No. 6,184,338.
- SUMM [0095] Various reactions may be carried out to produce acetylene-containing polymers. In these reactions, a sulfoxide is pyrolyzed to give the polyactylene and a sulfenic acid byproduct (RSOH) as shown in Formula II. For example, a benzenesulfenic acid may be eliminated from poly(phenyl vinyl sulfoxide)-containing copolymers to

produce polyacetylene-containing copolymers, such as poly(styrene-acetylene) block copolymers. Polyacetylene is typically difficult to work with because it is very insoluble in other materials. However, including it in a block structure allows the final structure to remain soluble.

- SUMM [0108] Diene, methacrylate and styrenic homo and block copolymers end-functionalized with alcohol(s), thiol(s), and/or amine(s) may be synthesized anionically by the use of suitable anionic initiators which contain protected functional groups that can be removed by post polymerization techniques. Suitable functional initiators are known in the art and are described in, e.g., U.S. Pat. No. 6,197,891, U.S. Pat. No. 6,184,338.
- SUMM [0119] The polymeric materials produced by acid-catalyzed pyrolysis of methacrylic and acrylic esters are methacrylic/acrylic acid or methacrylic/acrylic anhydride functionalized polymers. These acidanhydride-functionalized polymers may participate in further grafting reactions including esterification, amidation, and imidization reactions.
- SUMM [0121] In the case of amidation or imidization, the acid- or anhydride-functionalized polymeric material is subjected to reaction with amine nucleophiles. Suitable amines that participate in this reaction consist of, but are not limited to, typically primary alkyl, aryl, and alkylaryl-amines. The primary amines formula is RNH.sub.2 wherein R stands for an alkyl group having 1 to 10 carbon atoms, a cycloalkyl group having 5 to 12 carbon atoms, an aralkyl group having 7 to 22 carbon atoms or an aryl group having 6 to 12 carbon atoms.
- SUMM [0123] Diene, methacrylate and styrenic homo and block copolymers end-functionalized with alcohol(s), thiol(s), and/or amine(s) may be synthesized anionically by the use of suitable anionic initiators, which contain protected functional groups that can be removed by post polymerization techniques. Suitable anionic initiators are known in the art and are described in, e.g., U.S. Pat. No. 6,197,891, U.S. Pat. No. 6,160,054, U.S. Pat. No. 6,221,991, and U.S. Pat. No. 6,184,338. Objects and advantages of this invention are further illustrated by the following examples. The particular materials and amounts thereof, as well as other conditions and details, recited in these examples should not be used to unduly limit this invention.

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FILE 'USPATFULL, USPATOLD, USPAT2, CAPLUS, JAPIO' ENTERED AT 23:18:08 ON 08 JUN 2008

159 S (IMIDAT? OR IMIDIZ? OR IMIDIS?) AND SUPERCRITICAL?

L2 4630 S SMA (S) (POLYMER? OR COPOLYMER?)

L3 0 S L1 AND L2

L4 279533 S (STYREN? OR VINYL(2A)AROMATIC) (3A)COPOLY? OR POLY(1W)STYREN?(L5 11 S L1 AND L4 L5 ANSWER 10 OF 11 USPAT2 on STN

ACCESSION NUMBER: 2002:337815 USPAT2

Golf ball compositions including microcellular TITLE:

materials and methods for making same

INVENTOR(S): Harris, Kevin M., New Bedford, MA, United States

Rajagopalan, Murali, South Dartmouth, MA, United States

Cavallaro, Christopher, Lakeville, MA, United States PATENT ASSIGNEE(S): Acushnet Company, Fairhaven, MA, United States (U.S.

corporation)

NUMBER KIND DATE PATENT INFORMATION:

US 6676866 B2 20040113 US 2002-142168 20020510 20020510 (10) APPLICATION INFO.:

RELATED APPLN. INFO .: Division of Ser. No. US 2000-565108, filed on 4 May

2000, now patented, Pat. No. US 6386992

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED

FILE SEGMENT: GARAGE

FRIMARY EXAMINER: Kuhns, Allan R.

LEGAL REPRESENTATIVE: Swidler Berlin Shereff Friedman, LLP

NIMBER OF CLAIMS: 16

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 3 Drawing Figure(s); 1 Drawing Page(s) LINE COUNT: 973

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

1. A supercritical fluid (SCF) of an atmospheric gas is

injected into the polymer through a barrel to form a single-phase solution, with the SCF delivery system, screw, and injectors designed to allow for rapid dissolution;

DETD

Block polystyrene TPEs suitable for use in this invention include blocks of polystyrene or substituted polystyrene, e.g., poly(α-methyl styrene) or poly(4-methyl styrene) chemically linked or joined to the ends of lower softening point blocks of either an unsaturated or saturated rubber. Unsaturated rubber types typically include butadiene,

which can form styrene-butadiene-styrene (hereafter "SBS")

block copolymers, or isoprene, which can form styrene-isoprene-styrene (hereafter "SIS") block

copolymers, silicone rubber, balata, styrene-butadiene

rubber ("SBR"), and the like. Examples of suitable commercially available thermoplastic SBS or SIS copolymers include the KRATON® D series from Shell Corporation of Houston, Tex., which includes

KRATON® D2109, D5119 and D5298; VECTOR® from Dexco of Plaquemine, La.; and FINAPRENE® from Fina Oil and Chemical of Plano,

DETD Any conventional material or method may also be used in preparing the golf ball cover, which is typically disposed over the center or core. For example, as is well known in the art, ionomers, balata, and urethanes are all suitable golf ball cover materials. A variety of less conventional materials may also be used for the cover, e.g., thermoplastics such as ethylene- or propylene-based homopolymers and copolymers. These homopolymers and copolymers may also include functional monomers such as acrylic and methacrylic acid, fully or partially neutralized ionomers and their blends, methyl acrylate, methyl

methacrylate homopolymers and copolymers, imidized amino group-containing polymers, polycarbonate, reinforced polycarbonate, reinforced polyamides, polyphenylene oxide, high impact polystyrene, polyether ketone, polysulfone, poly(phenylene sulfide),

acrylonitrile-butadiene, acrylic-styrene-terephthalate, poly(ethylene terephthalate), poly(butylene terephthalate), poly(ethylene-vinyl

alcohol), poly(tetrafluoroethylene), and the like. Any of these polymers or copolymers may be further reinforced by blending with a wide range of density—adjusting fillers, including glass fibers or spheres, or metallic powders. The selection of a suitable cover, and application thereof over the mantle described herein, will be readily determinable by those of ordinary skill in the art, particularly in view of the disclosure herein.

=> log y COST IN U.S. DOLLARS	SINCE FILE	TOTAL SESSION
FULL ESTIMATED COST	91.21	91.84
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	-0.80	-0.80

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